Physiological characterization of stolon regression in a colonial hydroid

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Accepted 24 December 2007

SUMMARY
As with many colonial animals, hydactiniid hydroids display a range of morphological variation. Sheet-like forms exhibit feeding polyps close together with short connecting stolons, whereas runner-like forms have more distant polyps and longer connecting stolons. These morphological patterns are thought to derive from rates of stolon growth and polyp formation. Here, stolon regression is identified and characterized as a potential process underlying this variation. Typically, regression can be observed in a few stolons of a normally growing colony. For detailed studies, many stolons of a colony can be induced to regress by pharmacological manipulations of reactive oxygen species (e.g. hydrogen peroxide) or reactive nitrogen species (e.g. nitric oxide). The regression process begins with a cessation of gastrovascular flow to the distal part of the stolon. High levels of endogenous H2O2 and NO then accumulate in the regressing stolon. Remarkably, exogenous treatments with either H2O2 or an NO donor equivalently trigger endogenous formation of both H2O2 and NO. Cell death during regression is suggested by both morphological features, detected by transmission electron microscopy, and DNA fragmentation, detected by TUNEL. Stolon regression may occur when colonies detect environmental signals that favor continued growth in the same location rather than outward growth.

Key words: cell death, clonal organism, cnidarian, evolution and development, evolutionary morphology, hydroid, Podocoryna, Podocoryne, reactive oxygen species, reactive nitrogen species.

INTRODUCTION
As model systems for experimental studies of the evolution of development, clonal organisms (e.g. many fungi, herbaceous plants, and colonial invertebrates) are particularly useful because growth and development are inseparable, providing a broad chronological window in which manipulation is possible (Blackstone, 1997; van Kleunen and Fisher, 2001). In these clonal groups, the morphology can be idealized as comprising feeding and reproductive units, here termed polyps, which are interconnected by vascular stolons. Runner-like forms show widely spaced polyps and long stolonal connections, whereas sheet-like forms show closely packed polyps with short stolonal connections (Buss and Blackstone, 1991). These different morphological patterns may reflect changes in the timing of the production of polyps and stolon tips relative to rates of stolon growth and colony maturation; high ratios of production yield sheets, and low ratios yield runners.

Colonial hydroids can be particularly useful for investigating the differences between runners and sheets. These hydroids are representatives of cnidarians, a group of early-evolving animals. Considerable research has focused on the mechanisms underlying their development. Early studies concentrated on physiological parameters, particularly gastrovascular flow. For instance, Hale (Hale, 1964) described the growth of stolons in terms of the influence of gastrovascular flow on the cyclic pattern of forward surges and partial backward retraction in the stolon tip. Wytenbach (Wytenbach, 1968; Wytenbach, 1969; Wytenbach, 1973) further characterized this type of stolon elongation in terms of colony-level variations in growth rate, cycle phases, and features of gastrovascular flow. Belousov and collaborators (e.g. Belousov, 1973; Belousov et al., 1989) also studied stolon tip extension, but from the perspective of cell-level morphogenesis and physiology. Schierwater et al. (Schierwater et al., 1992) provided an overview of the various physiological parameters that contribute to gastrovascular flow and proposed a cellular mechanism for stolonal expansion and contraction. Dudgeon and Buss (Dudgeon and Buss, 1996) also emphasized the importance of gastrovascular flow in colony development. At the same time, a variety of studies focused on the effects of morphogens on colony growth (Plickert et al., 1987) including head activator (Schaller et al., 1989), stolon-inducing factor (Lange and Müller, 1991) and other peptides (Takahashi et al., 1997). More recent studies have begun to elucidate the role of gene activity in colony development. For instance, Cartwright and Buss (Cartwright and Buss, 1999) and Cartwright et al. (Cartwright et al., 1999; Cartwright et al., 2006) explored the expression of a well-studied parahox gene in the various components of the colony whose frequencies and distributions contribute to the overall pattern.

Despite these and other efforts to understand colony development, the relevance of regressive processes has not yet been considered in detail. For instance, stolon regression could influence both runner- and sheet-like patterns; the former could be the result of little regression of peripheral stolons, whereas the latter could be the result of greater amounts of such regression. Increasingly, developmental biologists are recognizing the widespread occurrence and importance of tissue regression and related processes and their contribution to pattern formation. For example, tissue regression plays a role in sculpting the individual digits of the limbs of vertebrates (Zuzarte-Luis and Hurlé, 2002) and the segments of the body of some invertebrates (Lohmann et al., 2002). The regression of the tadpole’s tail during its metamorphosis into a frog is another well-known example (Tata, 2006). Tissue regression is also important in the transition of insects from their larval forms to their adult forms and has been described in a variety of species (Hori et al., 2000).
Tissue regression may be relevant to pattern formation in colonial hydroids. In normally growing colonies, stolons can frequently grow out and then cease to grow or even under some circumstances regress. Stolon regression has been observed in untreated colonies on numerous occasions. While at any given time, only a few stolons in a colony may be regressing, over time such a process could have a cumulative effect on colony morphology. For example, consider two colonies, one with twice as many regressing stolons at any given time. Over weeks and months of colony development, the resulting difference in morphology could be substantial. Indeed, this change in morphology has been detected in colonies treated with vitamin C (Fig. 1). Nevertheless, the number of regressing stolons in colony morphology has been detected in colonies treated with vitamin C, though higher than controls, was relatively low and variable. Perhaps surprisingly, however, stolon regression was frequently observed to coincide with high levels of endogenous reactive oxygen species (ROS) production (Blackstone et al., 2004a; Blackstone et al., 2005). Thus, stolon regression may be an active process involving other active processes such as some form of regulated cell death. In this context, we developed a reliable method of pharmacologically inducing stolon regression in a colonial hydroid and have attempted to characterize this process at morphological, physiological and cellular levels.

MATERIALS AND METHODS
Study species and culture conditions
For all investigations, colonies of a single clone of Podocoryna (= Podocoryne) carnea Sars 1846 were cultured using standard methods (e.g., Blackstone, 1999). The clone, designated as P3, has been used in previous studies for over 14 years. All experiments were performed on clonal replicates grown from single polyp explants of a source colony. For each experiment, control and treated colonies were explanted from the same source colony to minimize non-genetic sources of variation (Ponczek and Blackstone, 2001). For measures of ROS, reactive nitrogen species (RNS), gastrovascular flow and cell death, colonies were grown on 15 mm diameter round glass coverslips. For transmission electron microscopy (TEM), clonal replicates were grown on a piece of clear polyethylene tied to a glass slide. All experiments were carried out at 20.5°C.

Treatment with hydrogen peroxide and nitric oxide
For characterization of stolon regression, it was deemed necessary to develop a protocol that reliably triggered the regression of a large number of stolons in a single colony. Since vitamin C-treated colonies typically show a variable response, colonies were treated with hydrogen peroxide (H$_2$O$_2$) or a nitric oxide donor, S-nitroso-N-acetyl-penicillamine (SNAP). For all H$_2$O$_2$ experiments, colonies were treated in a 5 mmol l$^{-1}$ solution of H$_2$O$_2$ and seawater (Vernole et al., 1998) with controls placed in plain seawater. For all NO experiments, colonies were treated in seawater with SNAP dissolved in dimethyl sulfoxide (DMSO) at a final concentration of 0.5 mmol l$^{-1}$. Controls were treated with an equivalent concentration of DMSO. For both H$_2$O$_2$ and NO experiments, a set of 6–7 treated and 6–7 control colonies were incubated for ~60 min beginning 24 h after feeding. Following treatment in respective solutions, the colonies were incubated in plain seawater and staining solution, typically for ~60 min prior to imaging or fixation. Excluding colonies analyzed for several of the cell death assays and for temporal sequence of events, fixation or imaging typically was done 2 h after treatment began.

Measurement of reactive oxygen species
The compound 2′,7′-dichlorodihydrofluorescein diacetate (H$_2$DCFDA; Molecular Probes, Eugene, OR, USA) was used as a semi-quantitative assay of general oxidative stress (Jantzen et al., 1998). This non-fluorescent dye is taken up by cells, and the acetate residues and clusters are among the primary mechanisms by which ROS can affect signaling pathways (Armstrong et al., 2004; Filomeni et al., 2005; Lee and Helmann, 2006; Salmeen et al., 2003; van Montfort et al., 2003). Thus, while there may be some debate as to whether the activation of H$_2$DCF is specific for the detection of H$_2$O$_2$ (Finkel, 2001), this assay quantifies the extent to which potential signaling pathways may be affected. A 10 mmol l$^{-1}$ stock solution of H$_2$DCFDA was prepared in anhydrous DMSO. After incubation in SNAP or H$_2$O$_2$, treatment solutions were replaced with plain seawater and H$_2$DCFDA was added to each dish to a concentration of 10 μmol l$^{-1}$. Colonies were then incubated in the dark for ~60 min. Subsequently, each colony was imaged in a microscope chamber containing plain seawater using an Orca-100 camera (Hamamatsu Photonics, Hamamatsu City, Japan) and an Axiovert 135 microscope (Carl Zeiss, Jena, Germany). ROS-related oxidative processes (as indicated by H$_2$DCFDA-derived fluorophores) were considered evident if statistically significant differences were detected compared to control colonies.
dichlorofluorescein) were imaged for three stolon tips per colony (excitation 450–490 nm, emission 515–565 nm). The luminescence and area of each stolon tip was measured in Image-Pro Plus (Media Cybernetics, Silver Spring, MD, USA) by: (1) using a bright-field image of the stolon tip to define the area of the stolon and the area of the background; (2) with these areas serving as ‘areas of interest’ for the fluorescent image, allowing the software to measure average luminescence of both areas; and (3) exporting these measurements to file. The area of each stolon was thus defined, and the luminescence of the stolon was adjusted for the background luminescence by subtraction. These measures were analyzed by a nested analysis of variance (stolon tips within clonal replicates, clonal replicates within treatments).

**Measurement of reactive nitrogen species**
The compound 4-amino-5-methylamino-2'7'-difluorofluorescein (DAF-FM) diacetate (Molecular Probes, Eugene, Oregon, USA) was used as a semi-quantitative indicator of low concentrations of NO (van Engeland et al., 1999; Wilson and Potten, 1999). After incubation of colonies in H2O2, treatment solutions were replaced with plain seawater and colonies were incubated for an additional ~60 min to mimic methods used for assaying ROS and RNS. Colonies (treated and controls) were then fixed and stained following the ApoDIRECT Assay protocol with slight modifications for adherent cells. Washing steps were performed in Petri dishes using 50 ml of wash buffer for each washing step. After the staining solution was prepared, the colonies were transferred to dry Petri dishes and the staining solution was pipetted onto the colonies. Rinsing steps were performed in Petri dishes using 50 ml of rinse buffer for each rinsing step. Following rinsing, the colonies were placed in 25 ml of propidium iodide (PI)/RNase staining buffer. Fragmented DNA was labeled using terminal deoxynucleotidyl transferase to catalyze the incorporation of fluorescein-12-dUTP at the free 3'-hydroxy-ends of the DNA. The stolon cells containing fragmented DNA were visualized using fluorescence microscopy (excitation 450–490 nm, emission 515–565 nm). All stolon cells (regardless of whether or not they contained fragmented DNA) were counterstained using the PI/RNase solution and were visualized using fluorescence microscopy (excitation 546 nm, emission >590 nm). Images were obtained, measured and analyzed as described above.

**Comparisons of stolon tip structure**
Stolons of control colonies and colonies treated with H2O2 were examined using transmission electron microscopy (TEM). Following an ~60 min treatment and a subsequent ~60 min incubation in plain seawater, colonies were fixed in 2.5% glutaraldehyde for 3 h at 4°C followed by three 10 min rinses in Millonig’s phosphate buffer. Specimens were postfixed in 1% osmium tetroxide for 2 h at room temperature followed by three 10 min rinses in Millonig’s phosphate buffer, then dehydrated in an ethanol series and cleared in acetone. Colonies were infiltrated and embedded in EMBed 812 resin (Electron Microscopy Sciences, Hatfield, PA, USA) and sectioned on a Reichert OmU2 ultramicrotome using a diatome diamond knife. Sections of approximately 90 nm in thickness were collected on formvar-coated slot grids or 75-mesh copper grids. Sections were stained with uranyl acetate for 20 min and with lead citrate for 40 min. Subsequently, sections were examined using a Hitachi H-600 transmission electron microscope. Micrographs of specimens were obtained using Kodak 4489 electron microscope film and negatives were then scanned.

**Comparisons of gastrovascular flow**
Videos of stolons lasting 10 min were taken at a point ~250 μm behind the tip of a stolon using the Axiovert microscope and a Dage MTI-72 camera. Three stolon tips per colony were videotaped. A total of five H2O2-treated colonies and five control colonies were treated ~30 min after feeding for ~60 min and were imaged in plain seawater 90–180 min after feeding and 10–100 min after treatment. Using Image-Pro Plus, 100 images were extracted from each video at 5 s intervals. Stolon width (perisarc to perisarc) and lumen width (endoderm to endoderm) were measured for each image. Measurements of maximum and minimum lumen width were obtained for three consecutive cycles. The period (in s) of each cycle was also measured. The rate of gastrovascular flow can be estimated as lumen width (maximum – minimum) divided by cycle period and stolon width (Blackstone, 1996). Although the two-dimensional analysis of flow requires certain assumptions (e.g. Bagatto and Bugggren, 2006), these assumptions seemed justifiable in this experiment.
Temporal sequence of stolon regression

To provide a better understanding of the temporal sequence of events during stolon regression, two sets of observations were carried out. First, an entire colony was treated with H$_2$O$_2$ for 1 h and observed using a dissecting microscope during and after treatment for >72 h. Second, eight stolon tips from eight different colonies were observed. Each colony was placed into a seawater-containing chamber for the Zeiss inverted microscope and videotaped prior to treatment for 5 min. H$_2$O$_2$ was then added to the chamber to a concentration of 5 mmol·l$^{-1}$. Colonies were continually taped until regression appeared to stop, which took anywhere from 30 to 90 min. Using Image-Pro Plus, images were extracted from each video at 30 s intervals. This time-lapse sequence was then examined and the timing of events associated with stolon regression was noted.

RESULTS

Measurement of reactive oxygen species and reactive nitrogen species

As found previously with vitamin C (Blackstone et al., 2004a; Blackstone et al., 2005), treatments with H$_2$O$_2$ and NO donors typically cause regression of peripheral stolons in $P$. carnea colonies as well as a flux of ROS and RNS (Figs 2, 3). Of 63 stolons measured from control colonies in the peroxide experiments, only one exhibited regression. Of 63 stolons measured from treated colonies in the peroxide experiments, 62 exhibited regression (comparing treated to controls, $\chi^2=118$, d.f.=1, $P<0.001$). At the treatment concentration (0.5 mmol·l$^{-1}$), fewer stolons regressed in the SNAP-treated colonies (28 out of 60), but still significantly more than the controls (3 out of 60; $\chi^2=27$, d.f.=1, $P<0.001$). When treated with either peroxide or SNAP (and whether visibly regressing or not), stolon tips exhibited high levels of endogenous peroxide-related fluorescence (Fig. 2, Fig. 4A,B). Those treated with exogenous H$_2$O$_2$ showed increased levels of ROS in stolon tips as compared to controls (Fig. 4A; $F=91.4$, d.f.=1, 12, $P<0.001$). Those treated with exogenous SNAP also showed increased levels of ROS in stolon tips as compared to controls (Fig. 4B; $F=14$, d.f.=1, 12, $P<0.01$). When treated with either peroxide or SNAP, stolon tips exhibited high levels of endogenous NO-related fluorescence (Fig. 3, Fig. 4C,D). Those treated with exogenous H$_2$O$_2$ showed increased levels of NO in stolon tips as compared to controls (Fig. 4C; $F=22.3$, d.f.=1, 12, $P<0.001$). Those treated with exogenous SNAP also showed increased levels of NO in stolon tips as compared to controls (Fig. 4D; $F=49.5$, d.f.=1, 10, $P<0.001$).

Note that in each experiment the probe for quantifying NO or ROS must enter a cell (and have acetate groups removed) before interacting with the target molecule and exhibiting fluorescence. Thus, although there does seem to be a pattern of higher fluorescence in the treatment that is also being detected, the probes are not simply reacting with the exogenous treatment. Given these results, the process of stolon regression appears to involve increased levels of both NO and ROS.

Fig. 2. Bright-field and fluorescent micrographs of stolon tips from colonies of $P$. carnea. (A,B) A healthy stolon tip (A) from a control colony exhibits relatively little peroxide-related fluorescence (B). (C,D) A regressing stolon tip (C) from a control colony exhibits an elevated level of peroxide-related fluorescence (D). (E,F) A regressing stolon tip (E) after 60 min treatment with 5 mmol·l$^{-1}$ H$_2$O$_2$ (F) exhibits an elevated level of peroxide-related fluorescence. (G,H) A regressing stolon tip (G) after 60 min treatment with 0.5 mmol·l$^{-1}$ of SNAP (H) exhibits an elevated level of peroxide-related fluorescence. Scale bar, 25 μm.

Fig. 3. Bright-field and fluorescent micrographs of stolon tips from colonies of $P$. carnea. (A,B) A healthy stolon tip (A) from a control colony exhibits relatively little NO-related fluorescence (B). (C,D) A regressing stolon tip (C) after 60 min treatment with 5 mmol·l$^{-1}$ H$_2$O$_2$ (D) exhibits an elevated level of NO-related fluorescence. (E,F) A regressing stolon tip (E) after 60 min treatment with 0.5 mmol·l$^{-1}$ SNAP (F) exhibits an elevated level of NO-related fluorescence. Scale bar, 25 μm.
Stolon regression in hydroids

Cell death analyses

Regressive processes in development frequently involve cell death. Many of the standard in vivo assays for cell death (Costa-Pereira and Cotter, 1999; Frey, 1995; van Engeland et al., 1999; Wilson and Potten, 1999) do not work well in the hydroid stolon system and yield variable and unreliable staining (data not shown). It is probable that the molecules involved in these assays do not reliably enter an intact stolon. To detect patterns of DNA fragmentation, TUNEL assays (in which stolons are fixed and permeabilized) did show a reliable pattern of staining (Fig. 5). In H2O2-treated colonies, the TUNEL fluorescent images showed a strong between-treatment difference (Fig. 6A; F=119, d.f.=1, 12, P<0.001), whereas the PI fluorescent images showed no difference between treatments (Fig. 6B; F=0.3, d.f.=1, 12, P>0.55). This suggests that there was no difference between treated and control stolons in the total number of cells (Fig. 5B,E), but at the same time there were many more cells with fragmented DNA in the treated stolons (Fig. 5C,F). In other studies, features of regulated cell death (e.g. DNA fragmentation, morphological changes, caspase activity) have been detected over a similar time course (Matsura et al., 2002; Blanco...
Comparisons of stolon tip structure
Ultrastructural features of healthy hydroid epitheliomuscular cells (EMCs) include a general columnar shape with one or more large vacuoles often surrounding the nucleus (Thomas and Edwards, 1991). Myoid processes, the contractile basal extensions of EMCs, are typically undetectable in stolon EMCs (Schierwater et al., 1992). Control and H2O2-treated stolons showed clear ultrastructural differences when visualized with TEM (Figs 7, 8). EMCs in control stolons had the characteristic columnar shape, typically with the cell nucleus suspended in a large vacuole (Fig. 7A–D) (Blackstone et al., 2004b), whereas the stolon tip maintained a smooth, rounded appearance (Fig. 7E). Treated stolons were often contracted (as indicated by the space between the ectoderm and the perisarc) and the cells showed extensive damage (Fig. 8A–E). The differences between cells of treated and control stolons cannot be explained entirely by contraction state of the stolon (e.g. the amount of stolon contraction in Fig. 7D and Fig. 8A appears approximately similar, but the cells look very different). Images show some recognizable features of cell death. Some of these features are characteristic of necrosis, such as flocculent condensation of chromatin (Fig. 8A,B) and vacuolation of the cytoplasm (Fig. 8A–E) (Syntichaki and Tavernarakis, 2003). Other features seem to be more characteristic of apoptosis, such as the lack of modification to cytoplasmic organelles such as mitochondria (Fig. 8C) (Syntichaki and Tavernarakis, 2003; Galluzzi et al., 2007). Compaction of the nucleus (Fig. 8A,B,D) and fragmentation of the plasma membrane into numerous small vesicles (Fig. 8A,E) also suggest apoptosis (Syntichaki and Tavernarakis, 2003; Galluzzi et al., 2007). Again, in other studies, features of regulated cell death (e.g. DNA fragmentation, morphological changes, caspase activity) have been detected over a similar time course (Matsura et al., 2002; Blanco et al., 2005; Mourdjeva et al., 2005). Not all of these features are found in all cells of treated stolons. However, none of these features has been found in any cells of non-regressing, control stolons. Indeed, Kroemer et al. (Kroemer et al., 2005) stress that it ‘must be remembered that dying in a cell population is not a synchronous but rather a stochastic process, and that at a given time, individual cells will be at different stages of the dying process.” Thus, according to the recommendations of the Nomenclature Committee on Cell Death, cells of treated stolons exhibited five of the eight morphological features that define apoptosis, one of the five features that define necrosis, and none of the features used to define autophagy (Kroemer et al., 2005).

Comparisons of gastrovascular flow
Colonies treated with H2O2 exhibit greatly diminished gastrovascular flow (Fig. 9). Using lumen width divided by cycle period and stolon width as a proxy for flow rate, analysis of variance indicates a highly significant difference between controls and treated colonies (F=115, d.f.=1, 8, P<0.001).

Temporal sequence of stolon regression
Along with stolon regression, other events occur during and after the treatment of a colony with H2O2. From whole-colony observation of a single treated colony (and cursory observation of numerous colonies), it appears that tentacles and polyps retract within 2 min of beginning the treatment. Shortly thereafter, the mouths of the polyps open and gastrovascular flow stops (within 5–15 min). Stolon regression was observed 15 min after treatment began. Some polyps were turning inside out after 20 min of treatment, but returned to normal configuration shortly thereafter. Within 1 h, the polyps were extended again while the tentacles remained retracted. After 1 h of treatment, the solution was replaced with plain seawater as part of the normal treatment protocol. Over the next 6 h, no change was detected in the colony except that the mouths of the polyps closed and more stolons regressed. All stolons had regressed within 4 h of treatment. The colony would not feed and gastrovascular flow could not be detected, despite the ability of the tentacles to stick to the shrimp and the ability of the polyps to move in response to the food source. The colony was able to feed again within 24 h following treatment. Gastrovascular flow was detected 2.5 h after feeding. Within 48 h after the initial treatment, regressed stolons had started to recover and the majority of stolons had recovered within 77 h. Notably, thinner stolon tips seem to regress before thicker stolon tips. Since thinner stolons typically receive less gastrovascular flow than thicker stolons (Dudgeon and Buss, 1996), this observation is consistent with the idea that gastrovascular flow is the proximate trigger of regression.

Close observation of stolon tips of eight colonies indicate that stolon regression follows a relatively stereotyped sequence of events. After treatment with H2O2, the first step of stolon regression involves the cessation of gastrovascular flow. In seven of the eight
colonies observed, this occurred within 0.5–2.0 min following treatment (13 min for the outlier). Direct observations of gastrovascular flow to tips suggest a somewhat quicker cessation of flow than whole-colony observations, possibly because (1) gastrovascular flow can be better observed at high magnification, or (2) flow to stolon tips ceases slightly before flow throughout the colony, or both. Shortly thereafter, the stolons begin to regress. In seven of the eight colonies observed, regression began 1.0–4.5 min after the initiation of treatment (49 min for the outlier). Finally, in seven of the eight colonies observed stolonal regression ceased within 8–22.5 min after treatment (90 min for the outlier). At this point, the stolon tip still appeared healthy. In some cases, regressing tips displayed a much more distorted appearance (e.g. Fig. 2G). This distortion featured a misshapen tip, often leaving small clusters of cells ‘stranded’ following regression (e.g. Fig. 2C). The almost immediate cessation of gastrovascular flow following treatment suggests that this event is the proximate trigger of stolon regression. Regression may thus be part of the regulation of a hydroid colony by ‘self inspection’ (Buss, 2001). Resolving the timing of the buildup of endogenous ROS and RNS relative to the cessation of gastrovascular flow proved difficult; for instance, the intense excitation wavelengths generated by the mercury-arc lamp of the Axiovert microscope may, by itself, affect gastrovascular flow to the stolon tip. Nevertheless, observations of numerous stolon tips (e.g. Figs 2 and 3) suggest that accumulation of endogenous ROS and RNS begins shortly after gastrovascular flow ceases.

**DISCUSSION**

As with many clonal organisms, colonial hydroids exhibit variation in growth form ranging from runner-like (widely spaced polyps and stolon branches) to sheet-like (closely spaced polyps and stolon branches). Rates of stolon branching relative to elongation have been thought to underlie this variation. We introduce the process of stolon regression as potentially relevant to this variation in growth form. Stolon regression occurs regularly in healthy colonies, and integrated over a long period of colony development even a modest rate of regression could potentially convert a runner-like colony into a sheet-like one. Higher rates of stolon regression may occur when colonies detect environmental signals that favor continued growth in the same location rather than outward growth. Previously, treatment with vitamin C has been shown to trigger stolon regression and sheet-like morphology (Blackstone et al., 2004a; Blackstone et al., 2005).
For detailed studies, many stolons of a colony can be reliably induced to regress by treatments with peroxide or nitric oxide. This process begins with a cessation of gastrovascular flow to the distal part of the stolon. High levels of endogenous ROS and RNS then accumulate as the stolon begins to regress. Remarkably, exogenous treatments with either H$_2$O$_2$ or a NO donor equivalently trigger endogenous formation of both H$_2$O$_2$ and NO. Transmission electron microscopy reveals that regressing stolons have atypical ultrastructural features. Cell death during regression is suggested by both TUNEL and TEM.

These results further support previous work suggesting a potentially large role for ROS and RNS in cnidarian signaling. Several examples will be briefly described to provide an appreciation of both the species and the mechanistic diversity of such signaling. First, Perez and Weis (Perez and Weis, 2006) suggest that symbiotic algae in heat-stressed anthozoans release ROS, which in turn trigger the release of NO from the host cells. By-products of this ROS–RNS signaling may contribute to anthozoan bleaching, that is, the release of the symbiotic algae. Second, Berking et al. (Berking et al., 2005) suggest that in some scyphozoans metabolically produced peroxides oxidize iodide to iodine. Endogenous tyrosine reacts with iodine to produce iodiferous tyrosine compounds. This process may trigger medusa formation, provide an oxidant defense system, and perhaps even supply the evolutionary roots of the vertebrate hormone thyroxin. Finally, building on earlier work, Doolen et al. (Doolen et al., 2007) suggest that ROS may have a crucial role in runner-like (moderate ROS) or sheet-like (low ROS) growth in P. carnea. The current research extends these findings by showing that, whereas moderate levels of ROS may lead to rapid colony growth and a runner-like form, high levels of ROS are involved in another process—stolon regression—which in turn may lead to more sheet-like growth. The taxonomic diversity of these results—in anthozoans, scyphozoans and hydrozoans—is matched by the diversity of pathways—host–symbiont interactions, medusa formation and colony development. The ‘many pathways’ view of ROS and RNS signaling in cnidarians is further supported by Blackstone et al. (Blackstone et al., 2005).

Fig. 8. TEM micrographs of a regressed stolon treated with 5 mmol l$^{-1}$ H$_2$O$_2$. (A–C) The epitheliomuscular cells (EMCs) in a regressed stolon exhibit vacuolation of the cytoplasm (A). Treated stolons are sometimes contracted (B; as indicated by the space between the perisarc and the ectoderm). Cells of a regressed stolon have relatively normal appearing mitochondria (C) and compact nuclei with flocculent condensation of chromatin (A,B). (D,E) Cells in the tip of the regressed stolon are shrunken (D) compared to their control counterparts (e.g. Fig. 7D). Once a stolon regresses, the cell membranes in the tip form numerous small vesicles (E). P, perisarc; E, ectoderm; MG, mesoglea; N, nucleus; M, mitochondrion; V, vacuole, Ve, vesicle. Scale bars, 1 $\mu$m.
Regulated cell death is increasingly accorded a prominent role in many biological processes, including those that involve tissue regression. Although programmed cell death is often equated with apoptosis, it is now clear that some forms of active or regulated cell death simply do not meet the criteria for apoptosis and that apoptosis is but one form of programmed or regulated cell death (e.g. Bredesen et al., 2006; Galluzzi et al., 2007). Thus, a plurality of cell death pathways and features is emerging (Sperandio et al., 2000; Proskuryakov et al., 2003; Syntichaki and Tavernarakis, 2003; Guimarães and Linden, 2004; Bredesen et al., 2006; Skulachev, 2006; Proskuryakov et al., 2003; Syntichaki and Tavernarakis, 2003; Blackstone, N. W. (1996). Gastrovascular flow and colony development in two colonial hydroids. Biol. Bull. Mar. Biol. Lab. Woods Hole 190, 56-58.


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